



# **LINKS and MS2LINKS USER MANUAL v. 1.0**

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# I. Introduction

## Overview

Structural elucidation of biomolecules and their assemblies is crucial in understanding the molecular basis of biological function. While high-resolution techniques like NMR and x-ray crystallography remain the premiere techniques, some biomolecular structures are still unknown, intractable to these techniques primarily because of challenges with size limitations and the dependence on crystal formation. To overcome these limitations, a growing number of alternative strategies that rely on sparse distance constraints (hydrogen/deuterium exchange (HDX),<sup>1, 2</sup> spin labeling,<sup>3</sup> fluorescence resonance energy transfer (FRET)<sup>4, 5</sup> have been employed to obtain structural information on such biomolecules.

In addition, mass spectrometric 3D (MS3D) approaches that couple chemical crosslinking and footprinting techniques with MS analysis has proven to be a valuable technique for the investigation of the 3D structure of proteins, nucleic acids, and macromolecular complexes.<sup>6-16</sup> The wide range and availability of crosslinking and footprinting reagents in conjunction with the flexibility of the MS analytical platform provides an excellent alternative technology for the structural elucidation of biomolecules that are not readily amenable to the traditional structural techniques. In this direction, new computational tools for the interpretation of mass spectra from crosslinked and modified proteins have been created.<sup>9, 17-22</sup>

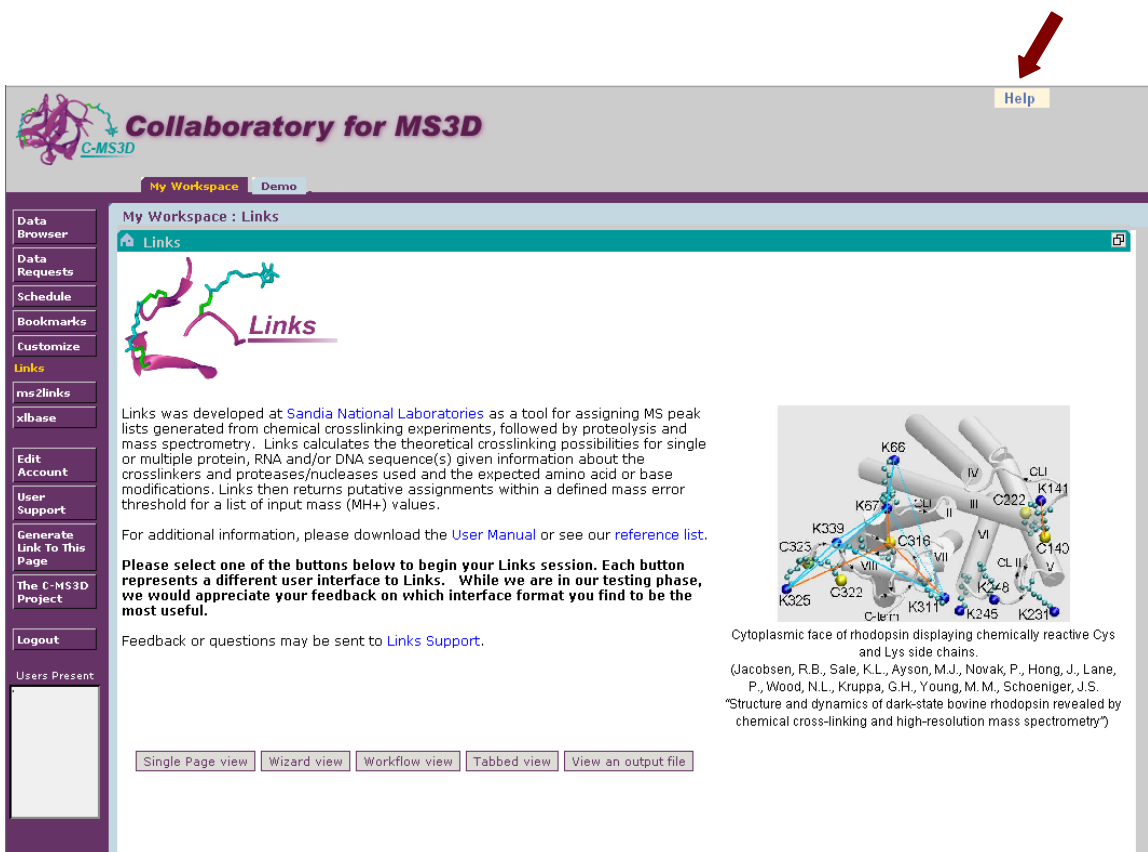
Links<sup>9,18</sup> was developed at [Sandia National Laboratories](#) as a program to analyze mass spectrometric data generated from native, modified, and crosslinked protein and nucleic acid digests. MS peak lists generated from chemical crosslinking or modification experiments, followed by proteolysis and mass spectrometry. Links calculates the theoretical crosslinking and modification possibilities for single or multiple protein, RNA and/or DNA sequence(s) given information about the crosslinkers and proteases/nucleases used and the expected amino acid or base modifications. Links then returns putative assignments within a defined mass error threshold for a list of input mass (MH+) values.

An analogous program, MS2Links, is used for assigning tandem MS peak lists generated from the fragmentation of crosslinked, modified or unmodified peptides, proteins and/or nucleic acids. MS2Links<sup>18, 19</sup> calculates the theoretical MS/MS fragment library given information about the identity of the base ion, crosslinkers (if applicable), desired ion types, and amino acid modifications. MS2Links then returns assignments within a defined mass error threshold for the list of input mass (MH+/MH-) values.

## Availability

As part of the C-MS3D (Collaboratory for MS3D) initiative, web-based implementations of these programs have been made available through our portal (<https://ms3d.ca.sandia.gov:11443/cms3d/portal>). To be able to access and use the programs, one needs to create an account in C-MS3D. Buttons that connect to the LINKS and MS2LINKS programs will appear at the left hand side of the webpage.

For general directions on how to navigate the portal, please refer to the portal manual. It can be accessed by clicking on the Help button at the upper right hand corner, then choose "How to".



The screenshot shows the 'Collaboratory for MS3D' web portal. At the top right, a red arrow points to a 'Help' button. The main content area is titled 'My Workspace : Links' and features a 'Links' section with a protein structure icon. Below this, there is a description of the Links tool, a link to the 'User Manual', and a list of buttons to begin a session. A sidebar on the left contains navigation links such as 'Data Browser', 'Data Requests', 'Schedule', 'Bookmarks', 'Customize', 'Links', 'ms2links', 'xlbase', 'Edit Account', 'User Support', 'Generate Link To This Page', 'The C-MS3D Project', 'Logout', and 'Users Present'. At the bottom, there are view options: 'Single Page view', 'Wizard view', 'Workflow view', 'Tabbed view', and 'View an output file'.

**Collaboratory for MS3D**

My Workspace : Links

**Links**

Links was developed at [Sandia National Laboratories](#) as a tool for assigning MS peak lists generated from chemical crosslinking experiments, followed by proteolysis and mass spectrometry. Links calculates the theoretical crosslinking possibilities for single or multiple protein, RNA and/or DNA sequence(s) given information about the crosslinkers and proteases/nucleases used and the expected amino acid or base modifications. Links then returns putative assignments within a defined mass error threshold for a list of input mass (MH+) values.

For additional information, please download the [User Manual](#) or see our [reference list](#).

Please select one of the buttons below to begin your Links session. Each button represents a different user interface to Links. While we are in our testing phase, we would appreciate your feedback on which interface format you find to be the most useful.

Feedback or questions may be sent to [Links Support](#).

Single Page view | Wizard view | Workflow view | Tabbed view | View an output file

Cytoplasmic face of rhodopsin displaying chemically reactive Cys and Lys side chains.  
(Jacobsen, R.B., Sale, K.L., Ayson, M.J., Novak, P., Hong, J., Lane, P., Wood, N.L., Kruppa, G.H., Young, M.M., Schoeniger, J.S.  
"Structure and dynamics of dark-state bovine rhodopsin revealed by chemical cross-linking and high-resolution mass spectrometry")

## II. Portal Structure

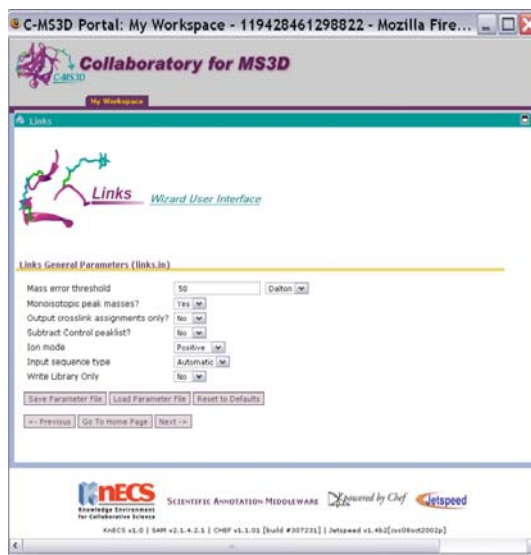
### General GUI

The steps in performing a Links session are standard, however different user interfaces are made available for Links and MS2Links, affording flexibility to users.

Single Page view



Wizard view



Workflow view



Tabbed view

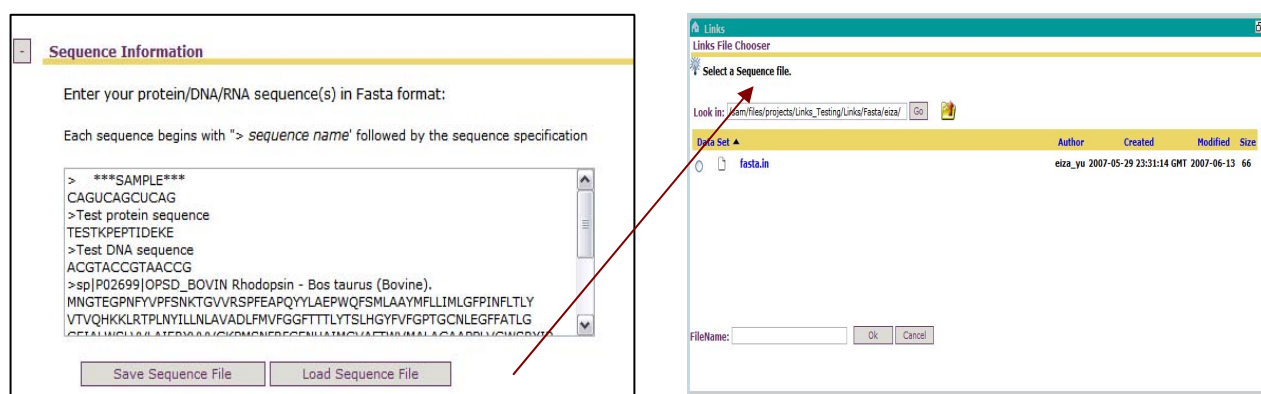



## Input/Output File structures and the Data Browser

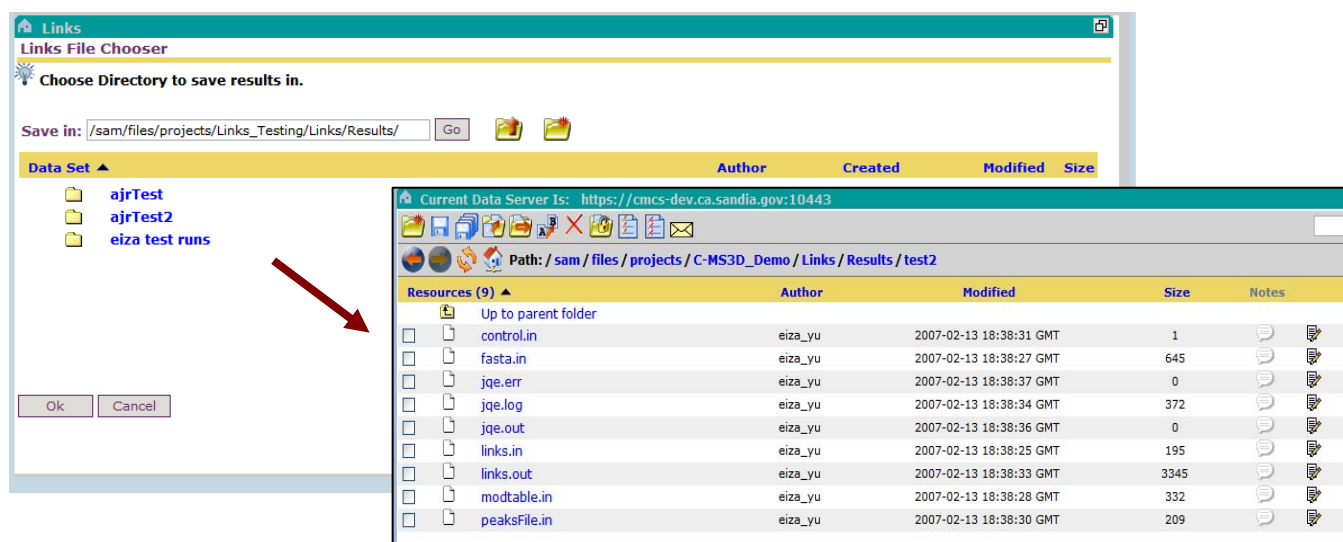
To run a Links job, the following input files are needed:

INPUT files	Description
fasta.in	Sequence/s of protein/nucleic acids to be analyzed
modtable.in	List of modifications/ cleavages to be applied in the analysis.
peaksFile.in	List of [M+H] <sup>+</sup> or [M-H] <sup>-</sup> observed in the mass spectra
control.in	List of [M+H] <sup>+</sup> or [M-H] <sup>-</sup> observed in control mass spectra (optional)
links.in	Defines analysis parameters

The input files can be saved by two ways: 1) copied in the input boxes and saved through the GUIs; or 2) uploaded and saved in the Links directory through the Data Browser. Each input file needs to be saved prior to running Links, otherwise you will get an error message. For more details on the input files, please refer to section III.



Once all necessary files are collected and saved, Links will run the job through the portal, and direct the user to the Data Browser, where all the output files and associated input files are saved under one folder in the user's own directory or team space directory. If one needs to control access to their files, permissions can be set through the  icon.



## Reporting Bugs

If you are having problems with Links/MS2Links, please refer to the FAQ section first to see if there is a quick fix to your problem. If it is not covered there or in this manual, please email us through the User Support Form.

## Citing C-MS3D portal

Yu,ET, Hawkins A, Kuntz ID, Rahn LA, Rothfuss A, Sale K, Young MM, Yang C, Pancerella CM and Fabris D. The Collaboratory for MS3D: A New Cyberinfrastructure Supporting the Structural Elucidation of Biological Macromolecules and their Assemblies Using Mass Spectrometry-based Approaches. J Proteome Res 2008, 7, (11), 4848-4857.

## Links and MS2Links were developed from the original ASAP and MS2Assign software:

Young MM, Tang N, Hempel JC, Oshiro CM, Taylor EW, Kuntz ID, Gibson BW, and Dollinger G. High throughput protein fold identification by using experimental constraints derived from intramolecular cross-links and mass spectrometry. Proc Natl Acad Sci USA 2000, 97, (11), 5802-6.

Schilling B, Row RH, Gibson BW, Guo X, and Young MM. MS2Assign, automated assignment and nomenclature of tandem mass spectra of chemically crosslinked peptides. J Am Soc Mass Spectrom 2003, 14, (8), 834-50.

Kellersberger KA, Yu E, Kruppa GH, Young MM, and Fabris D. Top-down characterization of nucleic acids modified by structural probes using high resolution tandem mass spectrometry and automated data interpretation. Anal Chem 2004, 76, (9), 2438-45.

### III. Running LINKS

#### 1. Sequence File

The target sequence can be entered directly in the sequence editor box using the standard one-letter code (use capital letters only), with each sequence starting with a sequence name (“>name”).

```
>testprotein
TESTKPEPTIDEKE
```

The input sequence can be saved as a \*.fasta file for future use. Alternatively, protein, DNA, and RNA sequences can be downloaded from databases in FASTA format, and loaded into the sequence editor box. When analyzing inter-molecular crosslinks, multiple sequences need to be entered into the sequence editor box and saved as a single fasta file. Links will automatically assign sequence numbers to the individual entries in the order that it appears in the file (See Modification Table).

#### 2. Modification Table

Links is capable of handling a variety of sequence modifications commonly encountered in protein and nucleic acid research. The user can define 1) amino acid-, 2) position-, or 3) peptide- specific modifications, as well as custom proteases/nucleases (XASES) and/or chemical crosslinkers, which are summarized and saved in a custom modification tables. Multiple modifications can also be defined in single session.

The Modification Table consists of the following sections:

1. TERMINAL MODIFICATIONS (applied to the termini of proteins and nucleic acids)
2. XASES (Protease,nuclease or custom cleavage applied to sequence)
3. NUCLEIC ACID MODIFICATIONS (applied to nucleic acid residues)
4. PROTEIN MODIFICATIONS (applied to amino acid residues)
5. CROSSLINKERS (generated from protein-protein or protein-nucleic acid crosslinking)

Links Modification Information (Xases, Modifications, Cross-linkers) (modtable.in)

Enter a modification table:

XASE *	RK	3	0	0	Tryp
MOD *	MW	15.9949	*	1	ox
MOD *	KX	226.0776	*	3	BT

Save Modification File

Load Modification File

Generate Modification Table

Reset to Defaults

<- Previous

Go To Home Page

Next ->

Fields:

Modifications - MOD\n" +  
(1) flag, (2) sequence number, (3) amino acid specificity string, (4) delta mass,  
(5) sequence position(s) to apply modification to (\*=all),  
(6) number of modifications allowed/peptide  
(7) description of modification

Protease/Nuclease - XASE  
(1) flag, (2) sequence number, (3) proteolytic specificity string,  
(4) Number of allowed missed cleavages, (5) N-terminal mass modification  
(6) C-terminal mass modification, (7) description

Crosslinker - XLINK  
(1) flag (2) sequence number (3) aa specificity string, site 1,  
(4) sequence number (5) aa specificity string,site 2  
(6) sequence position(s) for site 1 (7) sequence position for site 2  
(8) mass change upon cross-linking, (9) description



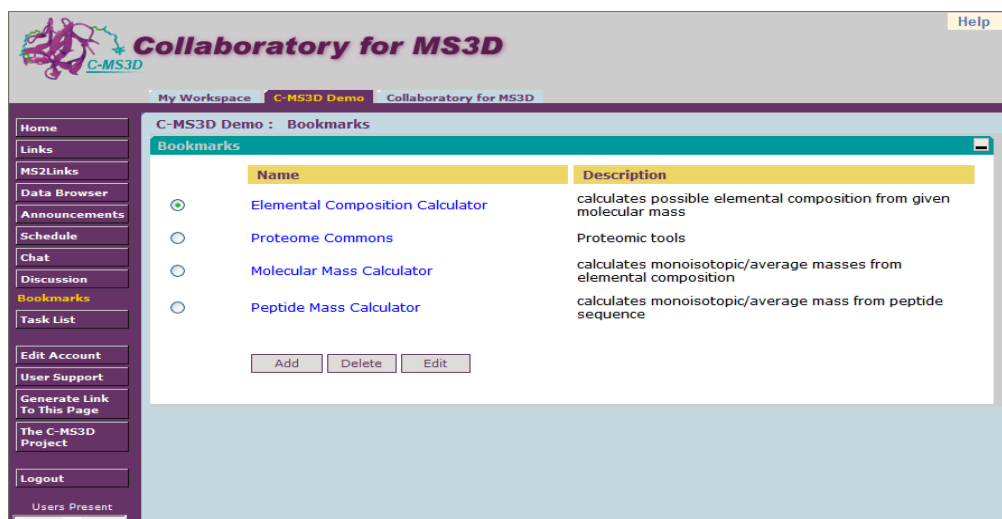
An easier way to generate the MOD Table is to use the Editor Form (click on the “Generate Modification Table” button). Modifications can be customized with the drop-down menu and the “Add Xase/Modification/Crosslinker” button. Once all the modifications are defined, the selection can either replace or added to the previous list of modifications.

The screenshot shows the 'Links Modification Table Editor Form' with three main sections: Xase(s), Modification, and Crosslinker. The Xase(s) section has a dropdown for 'Select Xase' set to 'User Defined' and a list of proteolytic specificities including Tryp, Chymotrypsin, V8 DE, Lys-C, Arg-C, Glu-C, Tyr-C, Asp-N, DE-N, pepsin, Carboxypeptidase, Aminopeptidase, Pronase, RNase T1, RNase T1 cyclic, RNase A, RNase A cyclic, and RNase T2. The Modification section has a dropdown for 'Select Modification' set to 'User Defined' and fields for 'Modification name' (OH-S'), 'Sequence Number (\* for all):' (1), 'Amino Acid specificity:' (X), 'Mass change (in Da):' (1.0078), 'Sequence position for modification (\* for all):' (\*), and 'Number of Modifications Allowed per peptide:' (\*). The Crosslinker section has a dropdown for 'Select Crosslinker' set to 'User Defined' and fields for 'Crosslinker Name' (DSS), 'Mass change (in Da):' (138.06809), 'Site' (1: 1, 2: 1), 'Amino Acid Specificity:' (KX), and 'Sequence Position (\* for all):' (\*). Red arrows point to the 'Add Xase', 'Add Modification', and 'Add Crosslinker' buttons. Below the form are buttons for 'Add to Modifications', 'Replace Modifications', and 'Cancel'.

Alternatively, the users can write their own modification table definitions with any text editor. Lines/entries in the modification table preceded by a # sign, indicates that these are comments or flags not in use. To define a modification necessary for a session, remove the # sign before the desired modification flag.

```
##### MODIFICATION TABLE#####
XASE *   RK| *   10   0       0   Tryp
#XASE *   DE| *   10   0       0   GluC
MOD *     MW    15.994914 *     *   ox-M
#MOD *     KX    226.077589 *     3   BTm
XLINK *     XK   *      XK   *     *   138.0681   DSS
MOD *     KX    156.0786 *     *   DSSOH
```

If you need to add custom modifications, refer to the following sections below for more details. If a modification is not available, one can create their custom Xase/Modification/Crosslinker using the User Defined selection, and simply change the values in the boxes to reflect the new values. To generate the needed monoisotopic and average mass shifts for the custom modification, we have provided links to peptide and molecular mass calculators through the Bookmarks tabs.



#### a. Defining terminal modifications

Protein and/or nucleic acid terminal modifications can be defined in END MODIFICATIONS.

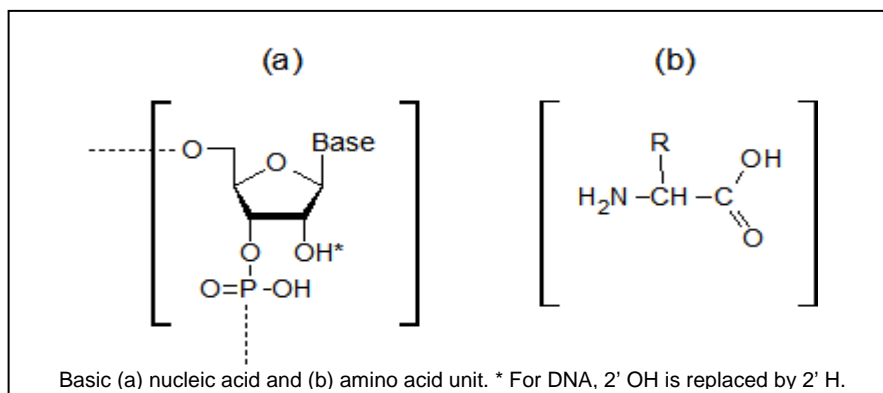
END MODIFICATION Fields:

- (1) Flag definition- MOD
- (2) Sequence number to which MOD will be applied to.  
To apply the MOD to all sequences, indicate (\*) or (all).
- (3) Sequence position to apply MOD  
X for N-term or 5'-end  
O for C-term or 3'-end
- (4) Delta mass to apply (refer to basic unit figure below)
- (5) Sequence position(s) to apply modification to (\*=all)
- (6) Number of modifications allowed per peptide (\*=defaults to 3)
- (7) A short description of the modification

```
##### END MODIFICATIONS #####
MOD 1 X 240.9067 * * PPP-5'
MOD 1 O -62.9635 * * OH-3'
```

In the example above, sequence 1 in the sequence.fasta input will have a triphosphate in the 5' end and a free hydroxyl in the 3' end. If the modifications are not provided in the list, the user can create any custom modification and add it in. To calculate delta mass (column 4), please refer to the figures below.

## PROTEIN AND NUCLEIC ACID BASIC UNITS



### b. Defining sequence cleavages

Protein and/or nucleic acid digestion/cleavages can be defined in XASES.

XASE Fields:

- (1) Flag definition- XASE
- (2) Sequence number to which XASE will be applied to.  
To apply the MOD to all sequences, indicate (\*) or (all).
- (3) Cleavage specificity string
- (4) Number of allowed missed cleavages
- (5) N-terminal/ 5'-end mass shift (after cleavage)
- (6) C-terminal/ 3'-end mass shift (after cleavage)
- (7) A short description of the modification

```
##### XASES#####
XASE 1 UC|* 10 1.0078 17.0027 RNase A
XASE 2 RK|* 3 0 0 Tryp
```

In the example above, sequence 1 in the fasta input will be digested with RNase A with a maximum of 5 missed cleavages and sequence 2 will be cleaved with trypsin (with a maximum of 3 missed cleavages).

The syntax to handle exceptions is the “^” (caret) symbol. To define trypsin cleavage rules with the exception (cleaves after K,R except if followed by a P on the C-term end):

```
##### XASES#####
XASE 2 RK|^P 3 0 0 Tryp
```

Users can specify multiple XASES to apply to 1 sequence (i.e. Trypsin/V8 combination) in a single Links session. The user can also create any custom XASE and add it to the list. To calculate associated mass shifts, please refer to the basic unit figures above.

⚠ When performing enzymatic digestion after modification or crosslinking will result to more missed cleavages and should be accounted for in the analysis. If a lot of missed cleavages are defined, LINKS will take more time to run the job. If possible, try to limit the allowed missed cleavages < 5.

### c. Defining base or amino acid residue modifications

Native and non-native modifications of proteins and nucleic acids can be defined in BASE or PROTEIN MODIFICATIONS.

MODIFICATION Fields:

- (1) Flag definition- MOD
- (2) Sequence number to which MOD will be applied.  
To apply the MOD to all sequences, indicate (\*) or (all).
- (3) Amino acid or base specificity string
- (4) Delta mass to apply (incremental mass shift due to modification)
- (5) Sequence position(s) to apply modification to (\*=all)
- (6) number of modifications allowed/peptide
- (7) A short description of the modification

```
##### BASE MODIFICATIONS #####  
MOD 2 STY 79.9663 52 1 PO4- (phosphorylation)  
MOD 2 K 42.0106 all 5 ace-K (acetylation)
```

In the example above, sequence 2 will have a one phosphorylation at position 52 and will look for all possible lysine acetylations. The user can specify multiple modifications in one session. Listed below are examples of other commonly encountered modifications.

```
#MOD 1 GCA 14.0156 * 3 DMS  
#MOD * A 73.0289 all 1 DEPC  
#MOD 1 M 15.9949 all 5 ox-M  
#MOD 1 ST 27.9949 all 5 formylation  
#MOD 1 XNKSTO 14.0156 all 5 met  
#MOD 1 Y 79.9568 all 5 sul- (sulfation)  
#MOD 1 Y 44.9851 all 5 nit- (nitration)  
#MOD 1 C 125.0477 all 10 nem-C (N-ethylmaleimide)
```

### d. Defining crosslinked substrates

Protein and/or nucleic acid crosslinking products can be defined in CROSSLINKERS.

CROSSLINKERS Fields:

- (1) Flag definition- XLINK
- (2) Sequence number to which XLINK will be applied.  
To apply the XLINK to all sequences, indicate (\*) or (all).
- (3) Amino acid specificity string for site 1
- (4) Sequence number
- (5) Amino acid specificity string for site 2
- (6) Sequence positions for site 1 (\*=all)
- (7) Sequence positions for site 2 (\*=all)
- (8) Mass change observed upon crosslinking
- (9) Description/Name of crosslinker

In the example below, Links will assign all possible DSS crosslinks between peptides generated from sequences in the fasta (In fact, Links can assign intra-molecular as well as inter-molecular crosslinked peptides, provided two sequences were entered in the fasta file and are defined in the parameter file(links.in)).

```
#####CROSSLINKERS#####
XLINK * KX * KX * * 138.06809 DSS
```

To limit the assignments to only inter-molecular crosslinks, one needs to specify the sequence numbers, as shown below as well as define it in the links parameter file (links.in):

```
XLINK 1 KX 2 KX * * 138.06809 DSS
```

In addition, the user can further limit the crosslinking assignments, if the sequence position for one of the sites is known. In this example, Links will assign GMBS crosslinks generated between C54 of protein 2 with all possible lysines in protein 1.

```
XLINK 1 KX 2 C * 54 165.04259 GMBS
```

In practice, one can assign type 0,1, and 2 crosslinks in a single MS by having the following mod table entries:

```
#####DSS CROSSLINKING MODIFICATIONS#####
XASE * RK|* 3 0 0 Tryp
MOD * KX 156.08 * 2 DSS-mono
XLINK 1 KX 2 KX * * 138.06809 DSS
```

⌘ The vast majority of user jobs still take just a few seconds to a couple of minutes actual compute time. The algorithms used in Links are exponential, meaning adding just a little bit more complexity greatly increases compute time, or inversely, efforts to simplify the problem a little bit may give very drastic increases in response time. There is no set time limit, but the server goes down for backups every Saturday so any running jobs are killed then.

Here are some ways to significantly reduce the number of combinations processed include:

1. Fixing the ends of the crosslinker:

Each XLINK sequence position that you can fix rather than leave as '\*' ought drop total combinations by a couple orders of magnitude.

2. Reducing the total number of allowed modifications for a MOD ought to halve the number of combinations.

Using a "\*" defaults to 3 applied modifications, so, putting in '2' or '1' instead will help.

3. Reducing the number of missed cleavages for the XASEs will also help, but the amount it helps is very dependent on where the XASE cleaves (anywhere from very little difference to orders of magnitude fewer combinations)

### 3. Input and Control Peaklist

Links needs a peaklist.txt file with the following format:

1	957.72	Y	1.851947e+07
1	964.01	Y	1.731260e+07
1	1000.71	Y	1.470965e+06
1	1034.74	Y	3.203310e+06
1	1056.04	Y	1.163770e+06
1	1153.88	Y	3.469220e+06
1	1169.91	Y	2.888720e+06
1	1199.87	Y	5.985100e+06
1	1213.5	Y	4.883800e+06
1	1213.5	?	1.126710e+07
1	1225.95	N	3.124410e+06

The first column is the spectrum/fraction number, second column is the reduced mass (M+H or M-H), the third column indicates whether input masses are considered monoisotopic, and the last column is the intensity. For example, "Y" indicates that the monoisotopic peak was observed in the spectra. "N" indicates that the monoisotopic peak was definitely not observed (thus the mass is the C13 peak in the spectra), and "?" indicates when the monoisotopic peak was not observed due to high baseline noise, this is especially true with clusters of higher charge states at high m/z values).<sup>18</sup> The significance of the 3rd column values will be discussed later in the links parameters section. More columns/data can be added to the peaklist file like charge, m/z, reduced monoisotopic mass, etc, however, the program only needs the minimum peaklist information shown above.

We have created translators to extract and reformat peaklists as links input files. The users only need to upload their peaklist into the portal. We currently support the following formats:

- \*.mzData
- \*.mgf
- \*.csv from Decon2LS (<http://ncrr.pnl.gov/software/Decon2LS.stm>)

#### Links Input Peak List (peaksFile.in)

Enter a peak list. The first column is the spectrum/fraction number, the second column is a MH+ or MH- value and the third column is Y, N or ?.

("Y" indicates that the mass is the first monoisotopic mass in a series, "N" indicates the mass is not, and "?" indicates that you're not sure.)

```
1 1270.70 Y
1 1277.07 Y
1 1285.65 Y
1 1296.71 Y
1 1312.71 Y
1 1324.75 Y
1 1327.65 Y
1 1359.62 Y
```

Save Peak List

Load Peak List

Load mzData Peak List

Load Decon2LS CSV Peak List

Load Mgf Peak List

Reset to Defaults

<- Previous

Go To Home Page

Next ->

For the peaklist translators, the user will be prompted for the following inputs. For mzData files, MS Level: 2 if data is from MS/MS experiments and 1 if data is from MS experiments. For mgf formats, the user will be asked to choose between average and monoisotopic mass. Lastly, for Decon2LS csv files, the user will be asked to choose between average and monoisotopic mass as well as positive or negative mode.

As a last resort, if the users' peaklist information is not saved in the above formats, one can modify their MH+/MH- peaklist using Excel to add the pertinent columns and save the output as .txt files.

Multiple peaklists (delineated by experiment #, in this case, 1-3) can be analyzed by merging peaklists in a single file:

1	957.72	Y	1.851947e+07
1	964.01	Y	1.731260e+07
1	1000.71	Y	1.470965e+06
2	1034.74	Y	3.203310e+06
2	1056.04	Y	1.163770e+06
2	1153.88	Y	3.469220e+06
3	1169.91	Y	2.888720e+06
3	1056.04	Y	1.163770e+06
3	1153.88	Y	3.469220e+06

The control peaklist has the same format and is optional. The user can input MS peaklists obtained from control reactions, which will then be subtracted from the input peaklist and will not be considered further in the assignment stage (differential MS analysis). Duplicate peaks will be subtracted from the input peaklist and will not be considered further in the assignment stage.

## 4. General Parameters (Links.in)

The Links.in file defines all the parameters needed for the automated assignment of digest peaks by Links.

**Links General Parameters (links.in)**

Mass error threshold: 50 PPM

Monoisotopic peak masses?: Yes

Output crosslink assignments only?: No

Subtract Control peaklist?: No

Ion mode: Positive

Input sequence type: Automatic

Write Library Only: No

Save Parameter File Load Parameter File Reset to Defaults

<- Previous Go To Home Page Next ->

### a. Mass error threshold:

Links will assign all possible matches for the experimental masses provided the values are within the maximum error threshold that the user has defined. The mass error can either be in parts per million (ppm) or in Daltons (Da). The accuracy in the calibration of the mass spectrum can guide the users as to what is an acceptable and practical error threshold value to define. Having a higher mass error threshold (~100 ppm) will make the search less stringent, but is necessary when the MS was obtained from lower resolution mass spectrometers.

If the peaklist contains MH<sup>+</sup> values with questionable monoisotopic peaks:

1 1213.5 ? 1.126710e+07

Links will try to match the experimental mass with both the theoretical monoisotopic (C12) mass as well as the C13 mass.

### b. Peaklist (Monoisotopic vs average mass list):

If the peaklist contains monoisotopic masses, select yes. This depends on the resolution of the user's mass spectrometer. Otherwise, selecting "No" means that the library of theoretical masses will be calculated and matched based on average mass.

### c. Output crosslink assignments only?

If the user is interested only in the assigned crosslinked species, then by selecting "Yes", Links will only write these assignments in the output.



#### d. Subtract Control peaklist?

Automated differential MS analysis can be performed when a control peaklist is available. Briefly, duplicate peaks will be subtracted from the input peaklist and will not be considered further in the assignment stage.

#### e. Ion Mode?

One can select between two modes (positive or negative) depending on which ion mode was used to obtain the mass spectra.

#### f. Input sequence type?

Links can recognize proteins, RNA, and DNA sequences automatically.

⌘ If dealing with RNA sequences without U residue, Links will automatically assign this sequence as DNA so the user needs to explicitly assign these sequences as RNA.

#### g. Write library only?

By selecting “Yes”, the user can generate the entire theoretical mass library that was defined with the given sequence file and modification table. Links writes out the theoretical mass list in the output and does not perform the matching/assignment sub-routine. All these parameter settings will be saved in a links.in file.

Links will be set to run a job once all the input files are defined, and the user will be prompted to enter a directory name for the folder where all associated files will be saved.



#### Links Job Submission

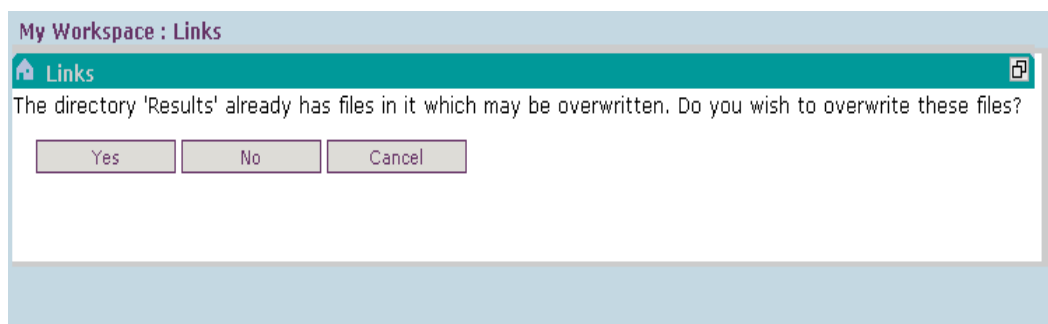
General Specified  
Sequence Specified  
Modifications Specified  
Peaks Specified  
Control Optional

Run Links Reset all

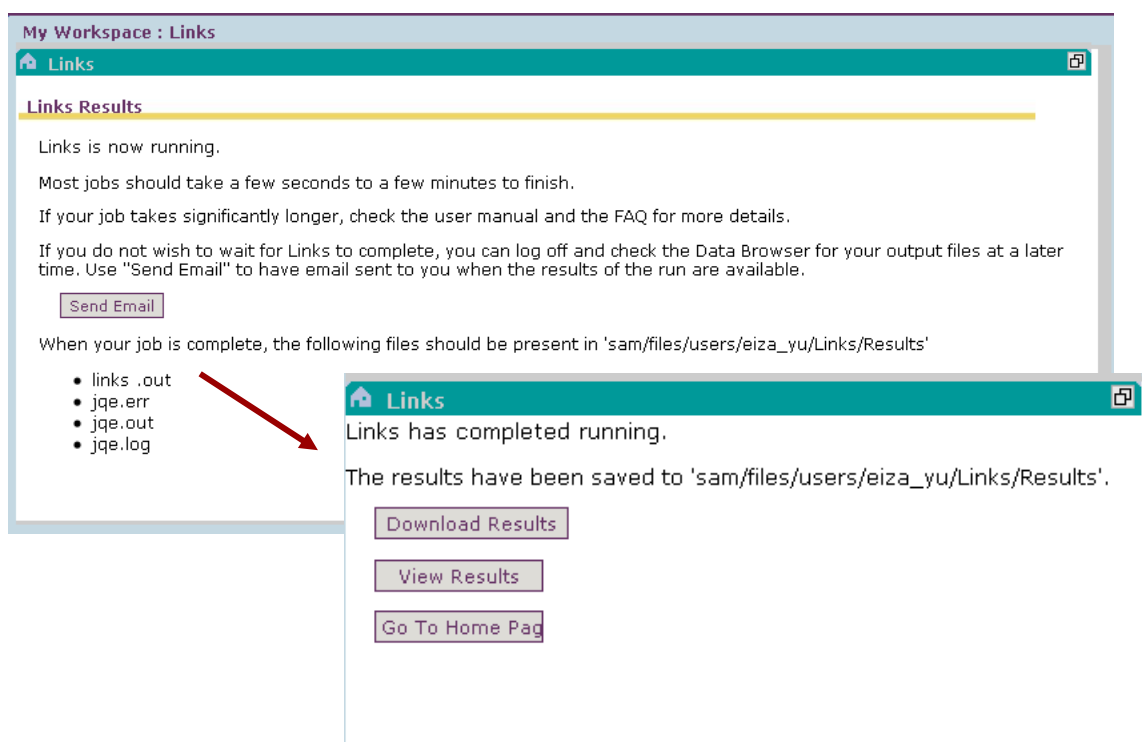
< - Previous Go To Home Page

A screenshot of the "Links File Chooser" dialog box. The window title is "My Workspace : Links". Inside, there's a "Links File Chooser" section with a "Choose Directory to save results in." label. Below this is a text field containing "/sam/files/users/test" and a "Go" button. To the right of the text field are two folder icons. Below the text field is a table with columns "Data Set", "Author", "Created", "Modified", and "Size". The table has one row with the data set "test", author "cms3d", created date "2007-11-05 17:36:02 GMT", modified date "2007-11-05", and size "0". At the bottom of the dialog are "Ok" and "Cancel" buttons. A red arrow points from the "Run Links" button in the "Links Job Submission" section to the "test" data set in the table.

If you try to overwrite an existing directory, you will be asked if you wish to proceed.



The page shown below will appear while Links is running. The output files will can be accessed from the Data Browser (please refer to the section on Output files). If the job takes longer than a couple of minutes to run, the user can log out of the portal (the job will not be terminated) and will be notified by email when the job is done.



## IV. Running MS2LINKS

MS2Links can be used to assign mass spectrum obtained from tandem mass spectrometry of entire proteins, nucleic acids, as well as digests. Similar to Links, MS2Links also has the ability to assign putative modifications (native, chemically modified, and crosslinks).

### 1. Sequence file

The sequence file requirements are similar to that of Links. The sequences of an entire protein, DNA or RNA can be inputs for MS2Links. For putative crosslinked peptide species (for example); the sequences of each peptide needs to be defined.

```
>testpeptide1
TESTKPEPTIDEKE

>testpeptide2
ACRSLINKPEP
```

### 2. Modification Table

MS2Links modification table also follows the same requirements as that of Links. For example, a putative crosslink between peptides in the sequence fasta file can be defined as:

```
#####DSS CROSSLINKING MODIFICATIONS#####
XLINK 1      KX      2      KX      *      *      138.06809      DSS
```

⌘ While the general parameters file takes care of the typical sequence ions generated from tandem MS, custom fragmentations[ref] due to a particular modification can be defined in the mod table. A well documented example, tandem MS of nucleic acids generally result to loss of nucleobases to form abasic sites and the presence of base modifications make this fragmentation more favorable.<sup>18, 23, 24</sup>

```
MOD  1      G  -151.049  *      1      -Gbase
```

Fragmentation of modified and/or crosslinked species do give rise to typical sequence ions, however, other fragmentation patterns seem to dominate the tandem MS. To get more details on the fragmentation behavior of such modified peptides, please refer to the following studies.<sup>25-31</sup>

```
MOD  1      K      140.084  *      *      DSSL13
MOD  1      K      224.165  *      *      DSSLint
```

### 3. Input and Control Peaklist

Please refer to the Links input and control peaklist section as the file requirements are similar.

### 4. General Parameters (MS2Links.in)

The MS2Links.in file defines all the parameters needed for the automated assignment of tandem MS peaks.

#### MS2Links General Parameters (ms2links.in)

Mass error threshold	<input type="text" value="100"/>	PPM <input type="button" value="v"/>
Monoisotopic peak masses?	Yes <input type="button" value="v"/>	
Assign peaklist, but only include crosslinks?	No <input type="button" value="v"/>	
Subtract control peaklist?	No <input type="button" value="v"/>	
Internal Ions	No <input type="button" value="v"/>	
Interfragment Crosslinks	No <input type="button" value="v"/>	
Input sequence type	Automatic <input type="button" value="v"/>	
<b>NOTE: The following two parameters will cause MS2Links to not make peak assignments.</b>		
Output theoretical library of crosslinked fragments?	No <input type="button" value="v"/>	
Output theoretical library of non-crosslinked fragments?	No <input type="button" value="v"/>	
A Ions	<input type="checkbox"/>	
B Ions	<input type="checkbox"/>	
C Ions	<input type="checkbox"/>	
D Ions	<input type="checkbox"/>	
W Ions	<input type="checkbox"/>	
X Ions	<input type="checkbox"/>	
Y Ions	<input type="checkbox"/>	
Z Ions	<input type="checkbox"/>	
<input type="button" value="Save Parameter File"/> <input type="button" value="Load Parameter File"/> <input type="button" value="Reset to Defaults"/>		
<input type="button" value="&lt;- Previous"/> <input type="button" value="Go To Home Page"/> <input type="button" value="Next -&gt;"/>		

#### a. Mass error threshold:

MS2Links will assign all possible matches for the experimental masses provided the values are within the maximum error threshold that the user has defined. The mass error can either be in parts per million (ppm) or in Daltons (Da).

b. Monoisotopic peak masses? (Monoisotopic vs average mass list):

If the peaklist contains monoisotopic masses, select yes. This depends on the resolution of the user's mass spectrometer. Otherwise, selecting "No" means that the library of theoretical masses will be calculated and will be matched based on average mass.

c. Assign peaklist, but only include crosslinks?

If the user is interested only in the assigned crosslinked species, then by selecting "Yes", MS2Links will only match the crosslinked species in the peaklist, and write these assignments in the output.

d. Subtract Control peaklist?

Automated differential MS analysis can be performed when a control peaklist is available. Briefly, duplicate peaks will be subtracted from the input peaklist and will not be considered further in the assignment stage.

e. Internal Ions?

MS2Links gives the user the option to include internal sequence ion arising from tandem MS.

f. Interfragment crosslinks

By selecting "Yes", ions generated from interfragment crosslinked species will also be considered.

g. Input sequence type?

MS2Links can recognize proteins, RNA, and DNA sequences automatically.

⌘ If dealing with RNA sequences without U residue, Links will automatically assign this sequence as DNA so the user needs to explicitly assign these sequences as RNA.

h. Output theoretical library of crosslinked fragments?

By selecting "Yes", the user will only generate the theoretical mass library of crosslinked fragments that was defined with the sequence files and the modification tables.

i. Output theoretical library of non-crosslinked fragments only?

By selecting "Yes", the user will only generate the theoretical mass library of non-crosslinked that was defined with the sequence files and the modification tables.

⌘ **NOTE:** By selecting "YES" on both the last two parameters (h and i), MS2Links will only write out the full theoretical mass list (as defined by the sequence and modification table) in the output and does not perform the matching/assignment.

#### j. Sequence Ion types

MS2Links gives the user the option to include all possible sequence ion types arising from tandem MS. For example, for low energy CID, the main ion types to consider are b- and y- ions. For more details, excellent reviews have been written about tandem MS of proteins<sup>30, 32</sup> and nucleic acids.<sup>33, 34</sup>

In addition to the ion types selected above, MS2Links will automatically assign water and NH<sub>3</sub> losses from fragment ions.

A Ions	<input type="checkbox"/>
B Ions	<input type="checkbox"/>
C Ions	<input type="checkbox"/>
D Ions	<input type="checkbox"/>
W Ions	<input type="checkbox"/>
X Ions	<input type="checkbox"/>
Y Ions	<input type="checkbox"/>
Z Ions	<input type="checkbox"/>

All these parameter settings will be saved in a ms2links.in file. Once all the input files are defined, MS2Links will prompt the user to enter a directory name for the folder where all associated files will be saved. Then, one can start the MS2Links session/job by clicking the “RUN MS2Links” button. The output files will can be accessed from the Data Browser (please refer to the section on Output files). If the job takes longer than a couple of minutes to run, the user can log out from the portal (the job will not be terminated) and will be notified by email when the job is done.

## V. Output files

Each Links/MS2Links session generates a directory in the C-MS3D portal Data Browser.

**Resources (4)**

	Author	Modified	Size	Notes
Up to parent folder				
Links	clyang	2006-04-12 21:42:13 GMT	0	
MS2Links	clyang	2006-04-12 21:42:20 GMT	0	
Open Babel files	clyang	2006-03-31 19:28:57 GMT	0	
UTR	eiza_yu	2007-01-18 05:51:30 GMT	0	

**Resources (6)**

	Author	Modified	Size	Notes
Up to parent folder				
Control	clyang	2006-04-12 21:42:13 GMT	0	
Fasta	clyang	2006-04-12 21:42:13 GMT	0	
Modifications	clyang	2006-04-12 21:42:13 GMT	0	
Parameters	clyang	2006-04-12 21:42:13 GMT	0	
Peaks	clyang	2006-04-12 21:42:13 GMT	0	
Results	clyang	2006-04-12 21:42:13 GMT	0	

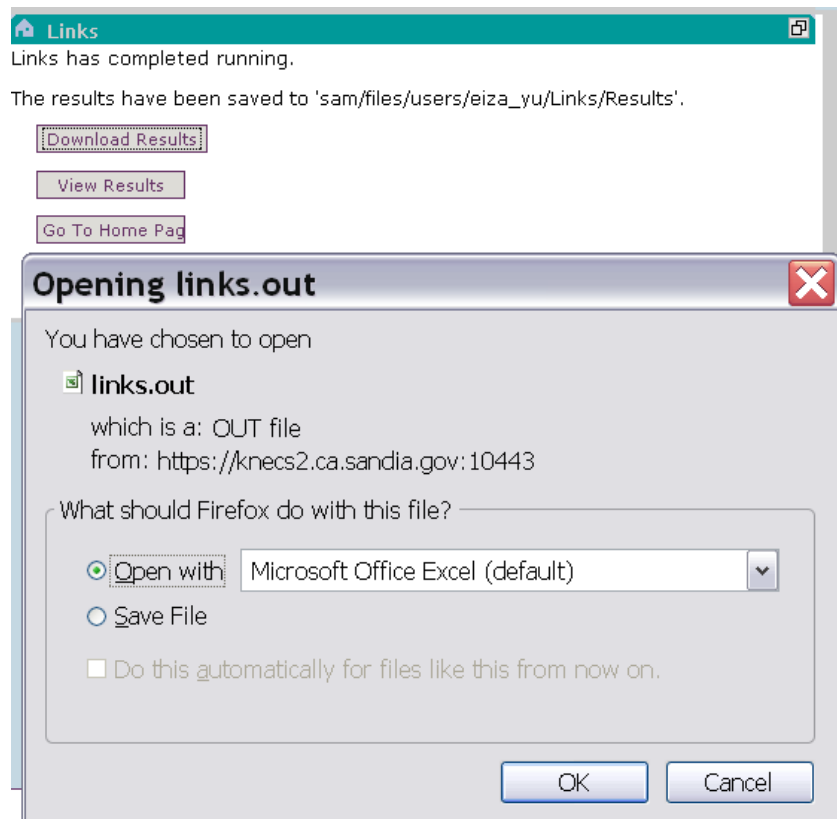
**Resources (9)**

	Author	Modified	Size	Notes
Up to parent folder				
control.in	clyang	2006-04-21 22:11:48 GMT	1	
fasta.in	clyang	2006-04-21 22:11:46 GMT	644	
jqe.err	clyang	2006-04-22 00:21:57 GMT	0	
jqe.log	clyang	2006-04-22 00:21:57 GMT	370	
jqe.out	clyang	2006-04-22 00:21:57 GMT	0	
links.in	clyang	2006-04-21 22:11:46 GMT	198	
links.out	clyang	2006-04-22 00:21:56 GMT	53864	
modtable.in	clyang	2006-04-21 22:11:46 GMT	4189	
peaksFile.in	clyang	2006-04-21 22:11:47 GMT	5141	

The directory folder contains copies of all the input files used in the Links session as well as runtime log and error files (jqe.\*). The jqe.\* files are logs of the actual links/ms2links program execution. So if something goes wrong, they can be used to help figure out what happened.

Resources (9) ▲	Author	Modified	Size	Notes
Up to parent folder				
control.in	clyang	2006-04-21 22:11:48 GMT	1	
fasta.in	clyang	2006-04-21 22:11:46 GMT	644	
jqe.err	clyang	2006-04-22 00:21:57 GMT	0	
jqe.log	clyang	2006-04-22 00:21:57 GMT	370	
jqe.out	clyang	2006-04-22 00:21:57 GMT	0	
links.in	clyang	2006-04-21 22:11:46 GMT	198	
links.out	clyang	2006-04-22 00:21:56 GMT	53864	
modtable.in	clyang	2006-04-21 22:11:46 GMT	4189	
peaksFile.in	clyang	2006-04-21 22:11:47 GMT	5141	

The user can also download the results directly to their computer by the “Download results now” button. A new pop-up window will appear and will prompt the user to either save or open the file in their workstation.





The resulting output file, links.out can be viewed within within the portal.

MS2Links

Output is only shown for matched peaks for this experiment  
For any additional result information, you must download the file,

[Download this file now](#)
[Go To Home Page](#)

Experiment Number	Sequence Number	MHobs	Desc	MHCalc	Error	Ion	Fragments
1	1	305.17	--	305.0166	0.2	d1+1OH-5'-H2O	U
1	2	305.17	--	306.0482	-0.9	y1+1p-3'	C
1	1	634.069	--	634.068	0.0	d2+1OH-5'-H2O	UA
1	1	979.116	--	979.1144	0.0	d3+1OH-5'-H2O	UAG
1	1	322.043	--	322.0431	-0.0	y1+1p-3'	C
1	1	322.043	--	323.0271	-1.0	d1+1OH-5'	U
1	1	667.09	--	667.0895	0.0	y2+1p-3'	GC
1	1	996.142	--	996.1409	0.0	y3+1p-3'	AGC
1	1	996.142	--	997.1249	-1.0	d3+1OH-5'	UAG
1	1	1303.0	--	1302.1653	0.8	MH(1)-1	UAGC
1	1	322.043	--	322.0431	-0.0	y1+1p-3'	C
1	1	322.043	--	323.0271	-1.0	d1+1OH-5'	U
1	1	651.095	--	650.5787	0.5	MH(1)-2	UAGC
1	1	651.095	--	652.0785	-1.0	d2+1OH-5'	UA
1	1	980.147	--	981.13	-1.0	c3+1OH-5'	UAG

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This viewer option only works if there were matched peaks in your spectra. Otherwise, you will get this message:

MS2Links

⚠ Could not find any matching peaks in the file

No matching peaks were found for this experiment.

For any additional result information, you must download the file,

[Download this file now](#)
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## The LINKS output file

In saved output files, the first part provides information on the defined parameters, sequence and modifications in the session:

```
Using MONOISOTOPIC masses.
Error is in PPM.
Reporting all assignments.
Amt of allowed error      : 50.000

LINKS: Automatic MS spectrum assignment program for cross-linked
macromolecules

Modifications:
MW      15.995      0      3      ox-M
KX      156.079     0      3      DSS-OH

Cross-link information: DSS(0-0)
      site1=XK
      site2=XK
      pos1=0
      pos2=0
      mass change = 138.068

Xase(s):
      Tryp (0)      : RK|*
      Mass mod<< 0.000
      Mass mod>> 0.000
      Missed cleavages: 5

2 modifications read in.

>APE
Length: 318
MPKRGKKGAVAEDGDELRTPEAKKSKTAAKNDKEAAGEGPALYEDPPD
QKTSPSAKPATLKICSWNVDGLRAWIKKKGLDWVKEEAPDILCLQETKCS
ENKLPaelQELPGLSHQYWSAPSDKEGYSGVGLLSRQCPLKVSYGIGDEE
HDQEGRVIVAEFDSFVLVTAYVPNAGRGLVRLEYRQRWDEAFRKFLKGLA
SRKPLVLCGDLNVAHEEIDLRNPKGNKKNAGFTPQERQGFGEllQAVPLA
DSFRHLYPNTPYAYTFWTYMMNARSKNVGWRLDYFLLSHSLLPALCDSKI
RSKALGSDHCPITLYLAL

Attempting to autodetect sequence type...

Input : protein sequence

Free memory4 sequences read into a 6348 member library...
```

The next part of the output file shows the assignments:

Expno	ExpMass	C13	Thr	Mass	Err	Seq #	Seq(s)	Int.	Sum	M.I.	Charge	Chrg, m/z
1	653.3089	--	0.0000	0.0	---							1.851947e+07 --
1	717.3847	--	717.3934	12.2	1	80-85				GLDWVK		1.470965e+06 --
1	823.3776	--	823.3738	4.6	1	188-193				WDEAFR		1.731260e+07 --
1	839.3725	--	839.3687	4.5	1	188-193 +ox				WDEAFRK		3.469220e+06 --
1	1137.5901	--	1137.5903	0.2	1	126-136				EGYSGVGLLSR		3.124410e+06 --
1	1403.7732	--	1403.8083	25.1	1-1	1-3, 195-202				MPK-FLKGLASR		4.883800e+06 --
1	1403.7732	--	1403.8262	37.7	1-2	26-31, 74-78				SKTAAK-AWI KK		5.985100e+06 --
1	2206.0691	--	2206.0259	19.6	1	86-103+1DSS-OH				EEAPDI LCLQETKCSENK		3.203310e+06 --
1	2719.4036(C13)		2718.3152	31.3	1-1	1-4, 86-103				MPKR-EEAPDI LCLQETKCSENK		5.565100e+06 -
.....												
Number of peaks assigned = 61/119 = 51.261 percent.												

The values for ExpNo, ExpMass, C13, Int.Sum, M.I., Charge, chrg, m/z columns are all copied from the input peaklist.in file( if available).

#### Some general explanations regarding the LINKS output file:

1. If the experimental mass is monoisotopic, the C13 column will show no value. However, in the case below the experimental mass was matched to the (C13) value of the theoretical mass.

Expno	ExpMass	C13	Thr	Mass	Err	Seq #	Seq(s)	Int.
1	2719.4036(C13)		2718.3152	31.3	1-1	1-4,86-103	MPKR-EEAPDILCLQETKCSENK	5.565100e+06

2. The sequence identifiers Seq# and Seq(s) provide the identity of the protein in the fasta file as well as the amino acid/nucleotide sequence of the assignment. For example; shows that the assigned peptide mass is from position 80-85 with sequence GLDWVK of sequence 1 in the fasta.in file.

Expno	ExpMass	C13	Thr	Mass	Err	Seq #	Seq(s)	Int.
1	717.3847	--	717.3934	12.2	1	80-85	GLDWVK	1.470965e+06 --

3. Crosslinked species are reported as (position#proteinA, position#proteinB) in the Seq(s) column. In the example below, the assignment is for an intra-molecular crosslink in protein 1, with position 1-3 crosslinked to 195-202.

Expno	ExpMass	C13	Thr	Mass	Err	Seq #	Seq(s)	Int.
1	1403.7732	--	1403.8083	25.1	1-1	1-3,195-202	MPK-FLKGLASR	4.883800e+06 --

The example below shows an assignment for an inter-molecular crosslink between position 26-31 of protein 1, and position 74-78 of protein 2.

Expno	ExpMass	C13	Thr Mass	Err	Seq #	Seq(s)	Int.
1	1403.7732	--	1403.8262	37.7	1-2	26-31,74-78 SKTAAK-AWIKK	5.985100e+06 --

4. Modified species are reported as (position# + MOD) in the Seq(s) column. In the example below, the assignment is for 2 biotinylated (BT) lysine residues in protein 1.

Expno	ExpMass	C13	Thr Mass	Err	Seq #	Seq(s)	Int.
1	1296.71	--	1296.674	27.6	1	1-7 + 2BT MPKRGKK	

This is also the same format for reporting intra-peptide crosslinks:

Expno	ExpMass	C13	Thr Mass	Err	Seq #	Seq(s)	Int.
1	2035.303	--	2035.33	13.4	1	45-50 + NM CGACUG	

## The MS2Links output file

The MS2Links output follows the same conventions as Links regarding crosslinks and modifications. As for the crosslinked sequence ions, MS2Links follows the same convention set in MS2Assign.<sup>19</sup>

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